PATENT COOPERATION TREATY

PCT

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY (Chapter I of the Patent Cooperation Treaty)

(PCT Rule 44bis)

Applicant's or agent's file reference 539.6000.10	FOR FURTHER ACTION	See item 4 below				
International application No. PCT/US2004/042792	International filing date (day/month/year) 17 December 2004 (17.12.2004)	Priority date (day/month/year) 17 December 2003 (17.12.2003)				
International Patent Classification (8th edition unless older edition indicated) See relevant information in Form PCT/ISA/237						
Applicant MEDTRONIC PHYSIO-CONTROL CORP.						

1.	This international preliminary report on patentability (Chapter I) is issued by the International Bureau on behalf of the International Searching Authority under Rule 44 <i>bis</i> .1(a).						
2.	This REPORT consists of a total of 8 sheets, including this cover sheet.						
	In the attached sheets, any reference to the written opinion of the International Searching Authority should be read as a reference to the international preliminary report on patentability (Chapter I) instead.						
3.	This report contains indications relating to the following items:						
	Box No. I	Basis of the report					
	Box No. II	Priority					
	Box No. III	Non-establishment of opinion with regard to novelty, inventive step and industrial applicability					
	Box No. IV	Lack of unity of invention					
	Box No. V	Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement					
	Box No. VI	Certain documents cited					
	Box No. VII	Certain defects in the international application					
	Box No. VIII	Certain observations on the international application					
4.	The International Bureau will communicate this report to designated Offices in accordance with Rules 44bis.3(c) and 93bis.1 but not, except where the applicant makes an express request under Article 23(2), before the expiration of 30 months from the priority date (Rule 44bis.2).						

	Date of issuance of this report 20 June 2006 (20.06.2006)
The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20, Switzerland	Authorized officer Athina Nickitas-Etienne
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Form PCT/IB/373 (January 2004)

PATENT COOPERATION TREATY

REC'D 2 1 JUL 2005

From the INTERNATIONAL SEARCHING AUTHORITY

To:				PCI			
see form PCT/ISA/220				WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY (PCT Rule 43 <i>bis</i> .1)			
				Date of mailing (day/month/year) see form PCT/ISA/210 (second sheet)			
	icant's or agent's file form PCT/ISA/22			FOR FURTHER ACTION See paragraph 2 below			
	International application No. International filing date PCT/US2004/042792 17.12.2004			day/month/year)	Priority date (day/month/year) 17.12.2003		
	national Patent Class N1/372, A61N1/3		both national classification A61N1/39	and IPC			
	icant DTRONIC PHYS	SIO-CONTROL	CORP.				
1.	This opinion co	ntains indicati	ons relating to the foll	owing items:			
	⊠ Box No. I	Basis of the op	oinion				
	☐ Box No. II	Priority					
	☑ Box No. III	Non-establishi	ment of opinion with rega	ard to novelty, inventiv	e step and industrial applicability		
	☑ Box No. IV	Lack of unity o					
	Box No. V Reasoned statement under Rule 43bis.1(a)(i) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement						
	☐ Box No. VI Certain documents cited						
	☐ Box No. VII Certain defects in the international application						
	☐ Box No. VIII Certain observations on the international application						
2.	FURTHER ACT	ION					
	If a demand for international preliminary examination is made, this opinion will usually be considered to be a written opinion of the International Preliminary Examining Authority ("IPEA"). However, this does not apply where the applicant chooses an Authority other than this one to be the IPEA and the chosen IPEA has notifed the						

International Bureau under Rule 66.1 bis(b) that written opinions of this International Searching Authority

If this opinion is, as provided above, considered to be a written opinion of the IPEA, the applicant is invited to submit to the IPEA a written reply together, where appropriate, with amendments, before the expiration of three months from the date of mailing of Form PCT/ISA/220 or before the expiration of 22 months from the priority date,

For further options, see Form PCT/ISA/220.

3. For further details, see notes to Form PCT/ISA/220.

Name and mailing address of the ISA:

will not be so considered.

whichever expires later.



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WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY

International application No. PCT/US2004/042792

	Вох	No	. <u>I</u>	Basis of the opinion
1.	With	reg ang	gard Juag	to the language , this opinion has been established on the basis of the international application in e in which it was filed, unless otherwise indicated under this item.
		land	auac	inion has been established on the basis of a translation from the original language into the following getuing yellowing the language of a translation furnished for the purposes of international search Rules 12.3 and 23.1(b)).
2.	With	req essa	gard ary to	to any nucleotide and/or amino acid sequence disclosed in the international application and on the claimed invention, this opinion has been established on the basis of:
	a. ty	pe ·	of m	aterial:
			a se	quence listing
]	table	e(s) related to the sequence listing
	b. fo	orma	at of	material:
	[in w	ritten format
			in c	omputer readable form
	c. ti	me	of fil	ing/furnishing:
	Į.		con	tained in the international application as filed.
	1]	filed	together with the international application in computer readable form.
	İ		furn	ished subsequently to this Authority for the purposes of search.
3		ha co	ıs be bies	tion, in the case that more than one version or copy of a sequence listing and/or table relating thereto ben filed or furnished, the required statements that the information in the subsequent or additional is identical to that in the application as filed or does not go beyond the application as filed, as oriate, were furnished.
4	Add	ditio	nal d	comments:

WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY

International application No. PCT/US2004/042792

	Box No. III Non-establishment of opinion with regard to novelty, inventive step and industrial applicability						
The obvi	The questions whether the claimed invention appears to be novel, to involve an inventive step (to be non obvious), or to be industrially applicable have not been examined in respect of:						
	the entire international application	on,					
\boxtimes	claims Nos. 7-19						
bec	ause:						
	the said international application, or the said claims Nos. relate to the following subject matter which does not require an international preliminary examination (specify):						
	the description, claims or drawi unclear that no meaningful opir	ngs <i>(</i> iion c	indicate particular elements below) or said claims Nos. are so could be formed (specify):				
	the claims, or said claims Nos. are so inadequately supported by the description that no meaningful opinion could be formed.						
\boxtimes	no international search report has been established for the whole application or for said claims Nos. 7-19						
	the nucleotide and/or amino acid sequence listing does not comply with the standard provided for in Annex C of the Administrative Instructions in that:						
	the written form		has not been furnished				
			does not comply with the standard				
	the computer readable form		has not been furnished				
			does not comply with the standard				
	the tables related to the nucleotide and/or amino acid sequence listing, if in computer readable form only, do not comply with the technical requirements provided for in Annex C-bis of the Administrative Instructions.						
	☐ See separate sheet for further details						

WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY

International application No. PCT/US2004/042792

	Во	x No. IV	Lack of unity of in	nvention						
1.	×	In resp	onse to the invitation	(Form P	CT/ISA/20	6) to pay addi	tional fees, the	applicant has	s:	
			paid additional fees.							
			paid additional fees	under pro	otest.					
		\boxtimes	not paid additional fe	es.						
2.		This Au	ithority found that the dicant to pay addition	requirer al fees.	nent of un	ity of invention	n is not complie	d with and cl	hose not to ir	vite
3.	Thi	s Author	ity considers that the	requiren	nent of uni	ity of inventior	ı in accordance	with Rule 13	3.1, 13.2 and	13.3 is
		complied	d with							
		not com	plied with for the follo	wing rea	sons:					
		see se	parate sheet							
4.	Co	nsequen	tly, this report has be	en estab	lished in r	espect of the	following parts	of the interna	ational applica	ation:
		all parts								
☑ the parts relating to claims Nos. 1-6										
	Bo	x No. V lustrial a	Reasoned statem applicability; citatio	ent und	er Rule 43 explanation	3 <i>bis</i> .1(a)(i) wi	ith regard to no	ovelty, inver	ntive step or	
1.	Sta	tement								
	No	velty (N)		Yes: No:	Claims Claims	1-6				
	lnv	entive st	ep (IS)	Yes: No:	Claims Claims	1-6				
	Ind	lustrial a	pplicability (IA)	Yes: No:	Claims Claims	1-6				
2.	Cit	ations a	nd explanations							

see separate sheet

Re Item IV.

The separate groups of inventions are:

Claims 1-6:

A patient parameter monitoring pod, comprising:

- a portable housing,
- a patient parameter module connectable to the patient through lead cables,
- a transceiver to communicate wirelessly to a defibrillator,
- and a data port to supply the patient data via a direct electrical connection to the defibrillator

Claims 7-12:

A patient parameter monitoring pod, comprising:

a housing holding a power supply;

patient lead cables attachable between the patient and the housing,

a carrying handle positioned to protect the patient lead cable port and the patient lead cables attached to the port from direct impact.

Claims 13-19:

A patient monitor pod system, comprising:

- a portable patient monitoring pod,
- a component bag,
- a patient parameter module,
- a data port,

wherein the component storage bag has pockets for holding the pod and components of the pod, the storage bag has openings exposing the data port and permits passage therethrough the patient lead cables.

They are not so linked as to form a single general inventive concept (Rule 13.1 PCT) for the following reasons: the common subject matter of the three groups of inventions is : a patient monitoring pod, comprising :

a housing.

patient lead cables attached between a patient and the housing to collect patient data, the

patient data including at least one vital sign.

These features are all disclosed in document US-A-5 105 821. For this reason, there is no unity between claims 1, 7 and 13.

Re Item V.

1 Reference is made to the following documents:

D1: EP 1 228 782 A (ST. JUDE MEDICAL AB) 7 August 2002 (2002-08-07)

D2: US 4 096 856 A (SMITH ET AL) 27 June 1978 (1978-06-27)

D3: US 5 105 821 A (REYES ET AL) 21 April 1992 (1992-04-21)

D4: EP 1 250 944 A (GE MEDICAL SYSTEMS INFORMATION TECHNOLOGIES,

INC) 23 October 2002 (2002-10-23)

2 INDEPENDENT CLAIM 1

The present application does not meet the criteria of Article 33(1) PCT, because the subject matter of **claim 1 does not involve an inventive step** in the sense of Article 33(3)PCT.

Document D3, which is considered to represent the most relevant state of the art to the subject matter of claim 1, discloses (the references in parentheses applying to this document): a patient parameter monitoring pod, comprising:

a **portable housing** (housing of element 14, figure 1) containing a power supply; a **patient parameter module** (element 14, figure 1) connectable to a patient via **lead cables** (leads connected to elements 39, figure 1) to collect patient data, the patient data including at least one vital sign;

and a data port (input connector 38, figure 1) adapted to supply the patient data via a direct electrical connection to the defibrillator (defibrillator 12, figure 1).

The subject-matter of independent claim 1 differs from the disclosure of D3 in that the patient parameter monitoring pod further comprises a **transceiver** adapted to

wirelessly transmit the patient data to a defibrillator.

The problem to be solved by the present invention may therefore be regarded as enabling the distance-communication between the pod and the defibrillator.

In view of D1 the solution proposed in claim 1 of the present application cannot be considered as involving an inventive step (Article 33(3) PCT) for the following reasons:

D1 discloses the same kind of apparatus of the one described in claim 1. In D1, the patient parameter monitoring pod (element 2, figure 1) comprises a transceiver (element 8, figure 1) adapted to wirelessly transmit the patient data to a defibrillator (element 4, figure 1).

Therefore the features disclosed in D1 and D3 would be combined by the skilled person, without exercise of any inventive skills in order to solve the problem posed. The proposed solution in independent claim 1 thus cannot be considered inventive (Article 33(3) PCT).

- Dependent claims 2-6 contain either features known per se from the prior art or being simple constructional features. Thus they would only satisfy Art. 33(2),(3) PCT when referring to a patentable independent claim.
- In order to facilitate the examination of the conformity of the amended application with the requirements of Art. 34(2)(b) PCT, the applicant is requested to **clearly identify the amendments carried out**, no matter whether they concern amendments by addition, replacement or deletion, and to indicate the passages of the application as filed on which these amendments are based (see also Rule 66.8(a) PCT).

If the applicant regards it as appropriate these indications could be submitted in handwritten form on a copy of the relevant parts of the application as filed.